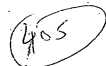
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SEARCH REQUEST FORM

Scientific and Technical Information Center

Requester's Full Name: PHTELSUDMAKER Examiner #: 77618 Date: 1 11/52	
Art Unit: 1/24 Phone Number 30 8 4 709 Serial Number: 59 8 3 9 2 8 9	
Mail Box and Bldg/Room Location: CM DE 17 Results Format Preferred (circle) PAPER DISK E-MAIL	
\mathcal{L}^{L}	
If m re than one search is submitted, please prioritize searches in order of need.	
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched.	
Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if	
KROWN Please allach a conviol the cover cheef, portinged plained and allaced	
STATHESIS & METHADS OF USE OF TETRAHYUKONJOLONE	
Title of Invention: ANALOGUES & DERIVATIVES	
Inventors (please provide full names): Point of Contact:	
DAYID B. FICK of Susan Hanley	
Farliest Priority Filing Date: 41 2 0 2 0 1	
CM1 12C14 Tel: 305-4052	
For Sequence Searches Only Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.	
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9 -N-(CH2)-(204	
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Searcher Phone #: 605-1155	AA Sequence (#)	Dialog			
Searcher Location:	Structure (#)		·		
Date Searcher Picked Up:	Bibliographic	Dr.Link	/~		
Date Completed: 1/16/02	Litigation	Lexis/Nexis	•		
Searcher Prep & Review Time:	Fulltext	Sequence Systems			
Clerical Prep Time:	Patent Family	WWW/Internet			
Online Time:	Other	Other (specify)			
PTO-1590 (8-01)		1.6.			

=> d all

CL28 ANSWER 1 OF 1 COPYRIGHT 2002 BEILSTEIN CDS MDLI

Beilstein Reg. No. (BRN): 1505538 Beilstein

Molecular Formula (MF): C15 H18 N2 O6

Autonom Name (AUN): methyl-carbamic acid 1-hydroxymethyl-5-methoxy-2,6-

dimethyl-4,7-dioxo-4,7-dihydro-1H-indol-3-ylmethyl

ester

Beilstein Reference (SO): 5-21 CAS Reg. No. (RN): 5904-30-3 Beilstein Pref. RN (BPR): 5904-30-3

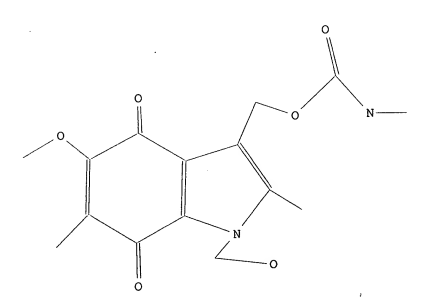
Formula Weight (FW): 322.32

Lawson Number (LN): 26215; 2817; 1762; 689; 289

Ring System Data:

Number of Rings (CNR): 2
Ring Systems (CNRS): 1
Diff. Ring Systems (CNDRS): 1
Ring Heteros (CNRH): 1
Acyclic Heteros (CNAH): 7

_	dex Ring System E (RF)	ormula BRIX Count
	-===+=================================	



Preparation: PRE

Reference(s):

1. Patent: Amer. Cyanamid Co., BE 653057 1963 Chem. Abstr., 64, <1966>, 15845e Melting Point:

Value | Ref. (MP) | (Cel) | 153.00 - 154.00 | 1

Reference(s):

 Patent: Amer. Cyanamid Co., BE 653057 1963 Chem. Abstr., 64, <1966>, 15845e

=> d all

CL24 ANSWER 1 OF 3 COPYRIGHT 2002 BEILSTEIN CDS MDLI

1553376 Beilstein Beilstein Reg. No. (BRN):

Molecular Formula (MF): C16 H20 N2 O6

1-(2-hydroxy-ethyl)-5-methoxy-2,6-dimethyl-3-Chemical Name (CN):

(methylcarbamoyloxy-methyl)-indole-4,7-dione

methyl-carbamic acid 1-(2-hydroxy-ethyl)-5-methoxy-2,6-dimethyl-4,7-dioxo-4,7-dihydro-1H-indol-3-

ylmethyl ester

5-21-13-00464 Beilstein Reference (SO):

10087-98-6 CAS Reg. No. (RN):

Beilstein Pref. RN (BPR): 10087-98-6 Formula Weight (FW): 336.34

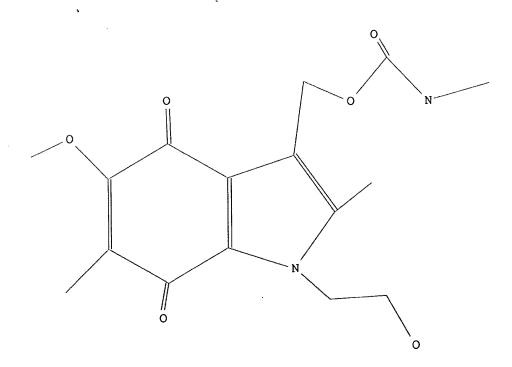
26215; 3122; 2817; 1762; 289 Lawson Number (LN):

Ring System Data:

Autonom Name (AUN):

Number of Rings (CNR): 1 Ring Systems (CNRS): Diff. Ring Systems (CNDRS): 1 Ring Heteros (CNRH): 1 Acyclic Heteros (CNAH):

Beilstein Ring (BRIX)		(RF)	Formula	•	BRIX Count
9.2.5-1.2-3.10	+======= 	===== C8N	 		1



Preparation:

PRE

Reference(s):

- 1. Allen; Weiss, J.Med.Chem., 10 <1967>, 23,29, CODEN: JMCMAR
- 2. Patent: American Cyanamid Co., US 3265698 1963 Chem. Abstr., 65, <1966>, 15330c

Melting Point:

Value (MP) (Cel)			Solv. (.SOL)	=======================================	Ref.
			•	petroleum	+==== 1
153.00	-	154.00	1		2

Reference(s):

- 1. Allen; Weiss, J.Med.Chem., 10 <1967>, 23,29, CODEN: JMCMAR
- 2. Patent: American Cyanamid Co., US 3265698 1963

Chem. Abstr., 65, <1966>, 15330c

Infrared Maximum:

IRM

Reference(s):

1. Allen; Weiss, J.Med.Chem., 10 <1967>, 23,29, CODEN: JMCMAR

Electronic Absorption Maximum:

EAM

Reference(s):

1. Allen; Weiss, J.Med.Chem., 10 <1967>, 23,29, CODEN: JMCMAR

CTUNCH Unchecked Data: NMR

Reference(s):

1. Allen; Weiss, J.Med.Chem., 10 <1967>, 23,29, CODEN: JMCMAR

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 L1
                   SCREEN 1839 AND 1994 AND 2004
 L2
                   SCREEN 2026 OR 2016 OR 2021 OR 1938
 L3
                 0 S L1 AND L2 NOT L3
 L4
       FILE 'REGISTRY' ENTERED AT 10:09:58 ON 16 JAN 2002
                 0 S L1 AND L2 NOT L3
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                   STR L1
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                   STR L1
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                   SAVE L13 SUD289/A
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       FILE 'HCAPLUS' ENTERED AT 11:53:51 ON 16 JAN 2002 do.

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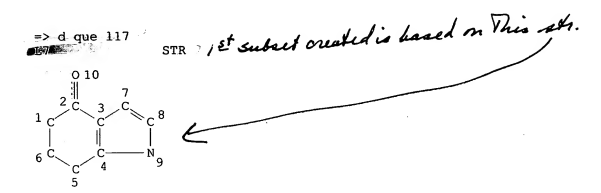
5 S L16 5 aits from L/6 aorapda.
< L17
-L18-
       FILE 'CAOLD' ENTERED AT 12:00:23 ON 16 JAN 2002
                0 S L15 no citations
 L19
       FILE 'BEILSTEIN' ENTERED AT 12:00:55 ON 16 JAN 2002

22 S L14 FULL 22 compds. Visor 114

19 S L20/COM 19 compdished
 L20
 L21
              3 S L21 NOT L22
3 S L23 AND PRE/FA Only 1st compd displayed due to high display cost
420 S WEISS?/AU AND ALLEN?/AU AND PY=1967
2 S L24 AND L25
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 (L24
 L25
                 2 S L24 AND L25
 L26
               1 S L27 AND PRÉ/FA one compd. disployed
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PATEL 09/839289

16/01/2002



NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

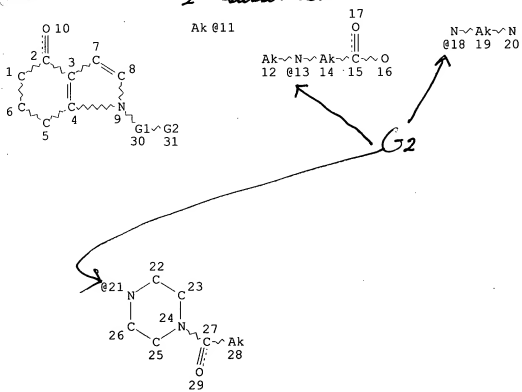
RSPEC I

NUMBER OF NODES IS 10

STEREO ATTRIBUTES: NONE

L9 1711 SEA FILE=REGISTRY SSS FUL L7

STR 2 nd subset sh.



VAR G1=11/CB VAR G2=OH/NH2/13/18/21 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED GRAPH ATTRIBUTES:

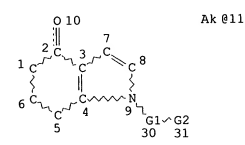
RSPEC I

NUMBER OF NODES IS 31

STEREO ATTRIBUTES: NONE

L13 29 SEA FILE=REGISTRY SUB=L9 SSS FUL L11

LIA STR Final subset search



17 0 N~Ak~N @18 19 20 Ak~N~Ak~C~O 12 @13 14 15 16 All Ak's are not substituted.

22 021 N C 23 | 24 | 27 26 C N 27 27 Ak 25 || 28

VAR G1=11/CB

VAR G2=OH/NH2/13/18/21

NODE ATTRIBUTES:

CONNECT IS E2 RC AT 11

CONNECT IS E1 RC AT 12

CONNECT IS E2 RC AT 14

CONNECT IS E2 RC AT 19

DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 31

STEREO ATTRIBUTES: NONE

L15 19 SEA FILE=REGISTRY SUB=L13 SSS FUL L14

L17 11 SEA FILE=HCAPLUS L15

PATEL 09/839289

16/01/2002

C=> d que 120
L14 STR

0 10 Ak @11 0 N~Ak~N

2 C 3 C 8 Ak~N~Ak~C~O
1 C C 8 12 @13 14 15 16

6 C C C N N

6 C C 3 G1~G2
5 30 31

VAR G1=11/CB
VAR G2=OH/NH2/13/18/21
NODE ATTRIBUTES:
CONNECT IS E2 RC AT 11
CONNECT IS E1 RC AT 12
CONNECT IS E2 RC AT 14
CONNECT IS E2 RC AT 19
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RSPEC I NUMBER OF NODES IS 31

STEREO ATTRIBUTES: NONE

L20 22 SEA FILE=BEILSTEIN SSS FUL L14

=> d ibib abs hitstr 1-11

L17 ANSWER 1 OF 11 HCAPLUS COPYRIGHT 2002 ACS 1997:479331 HCAPLUS ACCESSION NUMBER:

127:99527 DOCUMENT NUMBER:

Oxidative hair dye compositions containing TITLE: n-substituted 4-hydroxy indoline derivatives

Terranova, Eric; Fadli, Aziz; Lagrange, Alain INVENTOR(S):

Oreal S. A., Fr. PATENT ASSIGNEE(S):

Eur. Pat. Appl., 19 pp. SOURCE:

CODEN: EPXXDW Patent

DOCUMENT TYPE:

French LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 780118	A1	19970625	EP 1996-402297	19961029
EP 780118	B1	19971229		
R: DE, ES, E	R, GB	<u>, IT </u>		
FR 2742047	A1	19970613	FR 1995-14372	19951206
FR 2742047	B1	19980116		
CN 1189820	Α	19980805	CN 1996-195180	19960626
ES 2113769	Т3	19980501	ES 1996-402297	19961029
JP 09183716	A2	19970715	JP 1996-325758	19961205
JP 2996625	В2	20000111		
US 5755829	Α	19980526	US 1996-761756	19961205
US 6002018	Α	19991214	US 1998-14622	19980128
PRIORITY APPLN. INFO.:			FR 1995-14372 A	19951206
•			US 1996-761756 A3	19961205

MARPAT 127:99527 OTHER SOURCE(S):

Oxidative hair dye compns. contain n-substituted 4-hydroxy indoline derivs. (Markush structure given). A soln. of 8.85 g 4-hydroxy-1-N-(.beta.-hydroxyethyl)indole (prepn. given) in 40 mL of acetic acid was stirred with 1.9 g of sodium cyanoborohydride at 30.degree. for 30 min, then the reaction mixt. was poured into 200 g water at pH = 7.5 and the ppt. thus obtained was filtered and dried to obtain 7.75 g 4-hydroxy-1-N-(.beta.-hydroxyethyl)indoline (I). A hair dye prepn. contained I 0.895, paraphenylenediamine 0.540, water and excipient q.s. 100 g. The hair dye prepn. is mixed with equal amt. of 20 vol. hydrogen peroxide and applied to the hair.

186963-73-5P 186963-74-6P ΙT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (oxidative hair dye compns. contg. n-substituted 4-hydroxy indoline derivs.)

186963-73-5 HCAPLUS RN

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)- (9CI) (CA INDEX CN NAME)

RN 186963-74-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxypropyl)- (9CI) (CA INDEX NAME)

L17 ANSWER 2 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1997:168570 HCAPLUS

DOCUMENT NUMBER:

126:185981

TITLE:

Preparation of N-hydroxyalkyl-4-hydroxyindoles as

oxidative hair dye components

INVENTOR(S): Terranova, Eric; Fadli, Aziz; Lagrange, Alain

PATENT ASSIGNEE(S):

L'Oreal S. A., Fr.

SOURCE:

Eur. Pat. Appl., 20 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO. DATE
EP 754681 EP 754681	A1 B1	19970122 19980304	EP 1996-401413 19960626
		, ES, FR,	GB, IT, LI, NL, SE
FR 2736640	A1	19970117	FR 1995-8566 19950713
FR 2736640	В1	19970822	
CA 2222312	AA	19970130	
WO 9703049	A1	19970130	WO 1996-FR996 19960626
W: BR, CA,	CN, HU	, JP, KR,	MX, PL, RU
AT 163640	E	19980315	
ES 2117474	Т3	19980801	ES 1996-401413 19960626
JP 10512282	Т2	19981124	
BR 9609329	A	19990525	
JP 3095419	B2	20001003	JP 1997-505543 19960626
US 5704948	Α	19980106	US 1996-678981 19960712
US 5869692	Α	19990209	
PRIORITY APPLN. INFO).:		FR 1995-8566 A 19950713
			WO 1996-FR996 W 19960626
			US 1996-678981 A3 19960712

OTHER SOURCE(S):

MARPAT 126:185981

GI

Title compds. (I; R1 = hydroxyalkyl, alkoxyalkyl, aminoalkyl, etc.; R2, R3 AB = H, halo, alkyl, CO2H, alkoxycarbonyl, CHO; R4 = H, halo, alkyl, alkoxy, etc.) were prepd. Thus, 4-oxo-4,5,6,7-tetrahydrobenzofuran was cyclocondensed with H2NCH2CH2OH and the product dehydrogenated to give I (R1 = CH2CH2OH, R2-R4 = H). Data for activity of I were given.

186963-73-5P 186963-74-6P ΙT RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of N-hydroxyalkyl-4-hydroxyindoles as oxidative hair dye components)

RN 186963-73-5 HCAPLUS

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)- (9CI) CN

186963-74-6 HCAPLUS RN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxypropyl)- (9CI) (CA INDEX CN

$$\bigcap_{N} \bigcap_{\text{CH}_2-\text{CH}-\text{Me}}^{\text{OH}}$$

HCAPLUS COPYRIGHT 2002 ACS L17 ANSWER 3 OF 11 1989:594507 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

111:194507

TITLE:

Synthesis and oral hypoglycemic properties of

3-(1-oxo-3-hydroxy-2-cyclohexen-2-yl)-4-oxo-4,5,6,7-

tetrahydroindoles

AUTHOR(S):

Nagarajan, Kuppuswamy; Shenoy, Sharada J.; Talwalker,

Purnachandra K.

CORPORATE SOURCE:

Res. Cent., Hindustan Ciba-Geigy Ltd., Bombay, 400

063, India

SOURCE:

Indian J. Chem., Sect. B (1989), 28B(4), 326-32

CODEN: IJSBDB; ISSN: 0376-4699

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 111:194507

GT

AB Reaction of cyclohexane-1,3-diones with glyoxal, methylglyoxal, or phenylglyoxal afford bis -derivs. I (R, Rl = H, Me; R2 = H, Me, Ph) which are transformed to the title compds. by condensation with amines. Several of these are found to have activity in the fasted, glucose-primed rats. While many are as potent as tolbutamide, the activity of some is comparable to that of glybenclamide. Structure-activity relationships are discussed. Among these, cyclohexenylindole II has been chosen for further development.

IT 123271-86-3P 123271-89-6P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. and hypoglycemic activity of)

RN 123271-86-3 HCAPLUS

CN 4H-Indol-4-one, 1-(4-aminophenyl)-1,5,6,7-tetrahydro-3-(2-hydroxy-4,4-dimethyl-6-oxo-1-cyclohexen-1-yl)-2,6,6-trimethyl- (9CI) (CA INDEX NAME)

RN 123271-89-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-3-(2-hydroxy-4,4-dimethyl-6-oxo-1-cyclohexen-1-yl)-1-(4-hydroxyphenyl)-2,6,6-trimethyl- (9CI) (CA INDEX NAME)

HCAPLUS COPYRIGHT 2002 ACS ANSWER 4 OF 11

1989:457475 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

111:57475

TITLE:

Synthesis and oral hypoglycemic properties of

4-oxo-4,5,6,7-tetrahydroindole-3-acetic acids

AUTHOR (S):

Nagarajan, Kuppuswamy; Talwalker, Purnachand K.; Goud,

A. Nagana; Shah, Rashmi K.; Shenoy, Sharada J.; Desai,

Narasimha D.

CORPORATE SOURCE:

Res. Cent., Hindustan Ciba-Geigy Ltd., Bombay, 400

063, India

SOURCE:

Indian J. Chem., Sect. B (1988), 27B(12), 1113-23

CODEN: IJSBDB; ISSN: 0376-4699

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 111:57475

Condensation of .beta.-acetyl-2-hydroxy-4,4-dimethyl-6-oxo-1-cyclohexene-1-AΒ propionic acid (I) with NH4OAc and primary amines affords tetrahydroindole-3-acetic acids II (R = alkyl, aryl, aralkyl; R1 = Me, R2 = CH2CO2H), while another dimedone deriv. serves as starting material for isomeric indole-2-acetic acids II (R = alkyl, R1 = CH2CO2H, R2 = H).

4-Oxotetrahydroindole-2-carboxylic acids II (R = Ph, CH2CHMe2, R1 = CO2H, R2 = Me) and 3-carboxylic acids II (R = Ph, 4-FC6H4, R1 = H, R2 = CO2H), are obtained from the corresponding benzofurans. Some of the 3-carboxylic acid esters are transformed to tricyclic compds. like III [R3 = H, Me2N(CH2)3, Et2NCH2CH2]. Good oral hypoglycemic activity in normal rats is shown generally by the 3-acetic acids, among which C 8778-GO and C 9001-GO (II, R = Bu, CH2CHMe2, R1 = Me, R2 = CH2CO2H) are most active. These two acids are also active in streptozotocin-induced diabetic rats and have been investigated extensively. Structure-activity relationships are discussed.

IT 121626-09-3P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (prepn. and hypoglycemic activity of)

RN 121626-09-3 HCAPLUS

CN 1H-Indole-3-acetic acid, 4,5,6,7-tetrahydro-1-(4-hydroxyphenyl)-2,6,6-trimethyl-4-oxo- (9CI) (CA INDEX NAME)

IT 39991-82-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

RN 39991-82-7 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-phenyl- (9CI) (CA INDEX NAME)

L17 ANSWER 5 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1985:437321 HCAPLUS 103:37321

DOCUMENT NUMBER: TITLE:

Antiimplantation agents: part III -

1,2-diaryl-4,5-polymethylenepyrroles and

1,2-diaryl-4-oxo- and 1,2-diaryl-4-hydroxy-4,5,6,7-

tetrahydroindoles

AUTHOR (S):

Nagarajan, K.; Talwalker, P. K.; Shah, R. K.; Mehta,

S. R.; Nayak, G. V.

CORPORATE SOURCE:

Res. Cent., Hindustan CIBA-GEIGY Ltd., Bombay, 400

063, India

SOURCE:

Indian J. Chem., Sect. B (1985), 24B(1), 98-111

CODEN: IJSBDB; ISSN: 0376-4699

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 103:37321

GI

$$(CH_2)_n$$
 R
 $(CH_2)_n$
 R
 I
 $(CH_2)_n$
 R
 R
 R
 R

$$R^2$$
 R^2
 R^3
 R^4
 R^5
 R^5

Diarylpolymethylenepyrroles I (n = 1-3; R = Cl, OH, F, OMe; R1 = H, Cl) AB were prepd. by cyclocondensation of phenacylcycloalkanones II with RC6H4NH2. Alkylation of phenols I (R = OH) gave ethers I (R = OH) pyrrolidinoethoxy, piperidinoethoxy, morpholinoethoxy, Et2NCH2CH2O, etc.). Similarly prepd. were tetrahydroindoles III [R2, R3 = H, Me; R4 = H, Br, Cl, F, NO2, OMe, etc.; R5 = (substituted) Ph, pyridyl, HOCH2CH2, morpholinpropyl, chlorobenzyl, cyclohexyl; Z = 0], and some III (Z = 0) were reduced to give III (Z = H, OH). Several compds. exhibited antiimplantation activity in rats, among which the following were effective at a dose of 10 mg/kg orally for 6 days or less: ethers I [n =2, R = 4-Me2N(CH2)3O, 4-piperidinopropoxy, R1 = H; n = 3, R = 14-pyrrolidinoethoxy, R1 = H (IV)], oxoindoles III (R2 = Me; R3, R4 = H; R5 = 2-, 4-FC6H4, 4-pyrrolidinoethoxyphenyl, 4-NH2NHC6H4; Z = O), hydroxyindoles III [R2 = Me, R3, R4 = H, R5 = H, 4-FC6H4 (C 6924-Go); R2, R3 = Me, R4 = H, R5 = 4-FC6H4; Z = H, OH], and the deoxy deriv. III (R2-R5 = Me, H, H, 4-FC6H4, Z = H2) (V). Compds. IV and V (min. ED100 = 1 mg) and C 6924-Go (min. ED100 = 2 mg) showed no dissocn. between antiimplantation and estrogenic activities. Detailed studies on C 6924-Go showed that its activity is related to weak estrogenic-antiestrogenic properties. Structure-activity relationships were discussed.

39991-82-7P 68638-92-6P 96757-50-5P ΙT

96757-51-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and contraceptive activity of)

39991-82-7 HCAPLUS RN

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-phenyl- (9CI) CN INDEX NAME)

RN 68638-92-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(3-hydroxyphenyl)-6,6-dimethyl-2-phenyl- (9CI) (CA INDEX NAME)

RN 96757-50-5 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-(4-methoxyphenyl)-(9CI) (CA INDEX NAME)

RN 96757-51-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-(4-methoxyphenyl)-6,6-dimethyl- (9CI) (CA INDEX NAME)

IT 96757-31-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)

(prepn., alkylation, and contraceptive activity of)

RN 96757-31-2 HCAPLUS

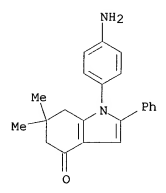
CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(4-hydroxyphenyl)-6,6-dimethyl-2phenyl- (9CI) (CA INDEX NAME)

IT 96757-34-5P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn., diazotization, and contraceptive activity of)

RN 96757-34-5 HCAPLUS

CN 4H-Indol-4-one, 1-(4-aminophenyl)-1,5,6,7-tetrahydro-6,6-dimethyl-2-phenyl-(9CI) (CA INDEX NAME)



L17 ANSWER 6 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

1980:75384 HCAPLUS

92:75384

TITLE:

Mass spectral fragmentations diagnostic of

1, 2-diaryl-6, 6-dimethyl-4-oxo-4, 5, 6, 7-

tetrahydroindoles

AUTHOR(S):

Ramadas, S. R.; Ramana, D. V.; Padmanabhan, S. Dep. Chem., Indian Inst. Technol., Madras, 600036,

India

SOURCE:

Indian J. Chem., Sect. B (1978), 16B(12), 1119-21

CODEN: IJSBDB; ISSN: 0376-4699

DOCUMENT TYPE:

CORPORATE SOURCE:

Journal

LANGUAGE:

English

AB The mol. ion (M+) peak is the base peak in all of the title spectra. Undergoes retro-Diels-Alder fragmentation, followed by extensive

rearrangement to give cyclic ions with concomitant loss of CO or at. H. The pyrrole moiety in M+ decomps. to aryl isocyanide ions.in another diagnostic path.

IT 68638-91-5 68638-92-6 68638-96-0

68638-99-3

RL: PRP (Properties)
 (mass spectrum of)

RN 68638-91-5 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyphenyl)-6,6-dimethyl-2-phenyl- (9CI) (CA INDEX NAME)

RN 68638-92-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(3-hydroxyphenyl)-6,6-dimethyl-2-phenyl- (9CI) (CA INDEX NAME)

RN 68638-96-0 HCAPLUS

CN 4H-Indol-4-one, 1-(2-aminophenyl)-1,5,6,7-tetrahydro-6,6-dimethyl-2-phenyl-(9CI) (CA INDEX NAME)

68638-99-3 HCAPLUS RN

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-6,6-dimethyl-2-CN phenyl- (9CI) (CA INDEX NAME)

L17 ANSWER 7 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1979:22730 HCAPLUS

DOCUMENT NUMBER:

90:22730

TITLE:

Studies on synthesis, chemical and spectroscopic

properties of 4-ketotetrahydroindole derivatives

AUTHOR(S):

Ramadas, S. R.; Padmanabhan, S.

CORPORATE SOURCE:

Dep. Chem., Indian Inst. Technol., Madras, India

SOURCE:

J. Prakt. Chem. (1978), 320(5), 863-72

CODEN: JPCEAO; ISSN: 0021-8383

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GT

$$\mathbb{R}^2$$
 \mathbb{N}^2
 \mathbb

- Oxotetrahydroindoles I (R1 = Ph, MeOC6H4, m- and p-NO2C6H4, o- and AB m-HOC6H4, and m- and p-HO2CC6H4, p-EtO2CC6H4, p-tolyl, o-H2NC6H4, benzyl, HOCH2CH2, 2-naphthyl, 6-quinolyl; R2 = H, Me) were prepd. in 62-95% yield by reaction of the phenacyldimedone II with R1NH2. The IR, UV and NMR data for I were tabulated.
- 68638-91-5P 68638-92-6P 68638-96-0P IT

68638-99-3P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (prepn. and spectra of)

68638-91-5 HCAPLUS RN

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyphenyl)-6,6-dimethyl-2-CN phenyl- (9CI) (CA INDEX NAME)

RN 68638-92-6 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(3-hydroxyphenyl)-6,6-dimethyl-2-phenyl- (9CI) (CA INDEX NAME)

RN 68638-96-0 HCAPLUS

CN 4H-Indol-4-one, 1-(2-aminophenyl)-1,5,6,7-tetrahydro-6,6-dimethyl-2-phenyl-(9CI) (CA INDEX NAME)

RN 68638-99-3 HCAPLUS

CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-6,6-dimethyl-2-phenyl- (9CI) (CA INDEX NAME)

L17 ANSWER 8 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1976:69246 HCAPLUS

DOCUMENT NUMBER:

84:69246

TITLE:

Carboxyarylindoles as nonsteroidal antiinflammatory

agents

AUTHOR(S):

Anderson, V. Brian; Agnew, Marc N.; Allen, Richard C.;

Wilker, Jeffrey C.; Lassman, Howard B.; Novick,

William J., Jr.

CORPORATE SOURCE:

Chem. Res. Dep., Hoechst-Roussel Pharm. Inc.,

Somerville, N. J., USA

SOURCE:

J. Med. Chem. (1976), 19(2), 318-25

CODEN: JMCMAR

DOCUMENT TYPE:

Journal English

LANGUAGE:

Fuditan

GI For diagram(s), see printed CA Issue.

Of 52 title compds., prepd. by condensation of an .alpha.-halo ketone with an enamine and cyclic condensation of the resulting 1,4-diketone with an aniline deriv., 34 had significant antiinflammatory activity and 5, including 3-(3-carboxy-4-hydroxyphenyl)-2-phenyl-4,5-dihydro-3H-benz[e]indole (I) [53597-27-6], were comparable to aspirin [50-78-2] in the carrageenin rat paw edema assay. Structure-activity relations were discussed.

IT 57859-78-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. and inflammation inhibiting activity of)

RN 57859-78-6 HCAPLUS

CN Benzoic acid, 2-hydroxy-5-(4,5,6,7-tetrahydro-4-oxo-2-phenyl-1H-indol-1-yl)- (9CI) (CA INDEX NAME)

L17 ANSWER 9 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1974:70733 HCAPLUS

DOCUMENT NUMBER:

80:70733

TITLE:

Reactivity of 4-oxo-4,5,6,7-tetrahydrobenzofurans. IV. Formation of 4,5-dihydropyrro[2,3-c]acridine

derivatives. Laboratory note

AUTHOR (S):

Takaqi, Kaname; Kobayashi, Noriaki; Ueda, Takeo

CORPORATE SOURCE:

Fac. Pharm., Univ. Kitasato, Tokyo, Japan

SOURCE:

Bull. Soc. Chim. Fr. (1973), (9-10, Pt. 2), 2807-9

CODEN: BSCFAS

DOCUMENT TYPE:

Journal

LANGUAGE:

French

GI

For diagram(s), see printed CA Issue.

The pyrroloacridines I (R = H, Me, Et, CH2CH2OH, Ph, 2-naphthyl, AB C6H4OMe-p; R1 = Me, Ph) were prepd. by cyclizing 2-benzoylmethyl-4,5dihydroresorcinol with RNH2 to give the indolones II (X = 0), which were reduced with N2H4 to II (X = H2) and cyclized with o-R1COC6H4NH2.

39991-82-7P IT

RL: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

39991-82-7 HCAPLUS RN

(CA 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-phenyl- (9CI) CN

INDEX NAME)

L17 ANSWER 10 OF 11 HCAPLUS COPYRIGHT 2002 ACS

ACCESSION NUMBER:

1973:71906 HCAPLUS

DOCUMENT NUMBER:

78:71906

TITLE:

Substituted 2-phenyl-4,5,6,7-tetrahydroindoles

INVENTOR(S):

Luecke, Bernhard; Lehman, Gerhard

SOURCE:

Ger. (East), 3 pp.

CODEN: GEXXA8

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DD 86826		19720105	DD 1970-145212	19700130

For diagram(s), see printed CA Issue. GI

Tetrahydroindolones I (R = H, OH; R1 = H, OMe) were prepd. in 52-84% yield AB by treating the corresponding 2-phenacyl-1,3-cyclohexanedione with RCH2CH2NH2. I (R = OH, R1 = H) was reduced with NaBH4 to give 78% of the 4-hydroxy deriv.

TT 39991-82-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

39991-82-7 HCAPLUS

RN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(2-hydroxyethyl)-2-phenyl- (9CI) CN INDEX NAME)

HCAPLUS COPYRIGHT 2002 ACS L17 ANSWER 11 OF 11

1972:434240 HCAPLUS ACCESSION NUMBER:

77:34240 DOCUMENT NUMBER:

Reaction of pyrrole ketones with formaldehyde. TITLE:

Formation of N-pyrrolemethanols

Berger, Joel G.; Schoen, Karl AUTHOR(S):

Endo Lab., Inc., Garden City, N. Y., USA CORPORATE SOURCE:

J. Heterocycl. Chem. (1972), 9(2), 419-21 SOURCE:

CODEN: JHTCAD

DOCUMENT TYPE: Journal

English LANGUAGE:

For diagram(s), see printed CA Issue. GΙ

N-Pyrrolemethanols are prepd. by heating 3-acylpyrroles with formaldehyde AB in the presence of NaOH. 3-Acetyl-2,5-dimethylpyrrole (I) is converted to the corresponding N-pyrrolemethanol (II), while 2,3-disubstituted 4-oxo-4,5,6,7-tetrahydroindole-1-methanols (III) are obtained from the corresponding indoles (IV). 3 III, which can contain a 2-Me and a 3-Me or 3-Et group are prepd. Similarly prepd. is 2-ethyl-3-methyl-6-oxo-4,5dihydro-6H - cyclopenta[b]pyrrole - 1-methanol.

36764-23-5P 36784-83-5P 36827-21-1P ΙT

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of)

36764-23-5 HCAPLUS RN

4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(hydroxymethyl)-3-methyl- (9CI) (CA CN

36784-83-5 HCAPLUS RN

4H-Indol-4-one, 3-ethyl-1,5,6,7-tetrahydro-1-(hydroxymethyl)-2-methyl-CN (9CI) (CA INDEX NAME)

RN 36827-21-1 HCAPLUS CN 4H-Indol-4-one, 1,5,6,7-tetrahydro-1-(hydroxymethyl)-2-methyl- (9CI) (CA INDEX NAME)

=> d ibib abs hitstr

L18 ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:762968 HCAPLUS

DOCUMENT NUMBER: 135:304105

TITLE: Preparation of nucleosides and isoindolinone

derivatives as anti-inflammatory agents

INVENTOR(S): Japtap, Prakash; Southan, Garry; Salzman, Andrew;

Szabo, Csaba; Ram, Siya
PATENT ASSIGNEE(S): Inotek Corporation, USA
SOURCE: PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

Pi	ATENT	NO.		KI	ND	DATE			A.	PPLI	CATI	ои ис	0.	DATE			
W	2001													2001			
	W:-	AE.	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	_CN,_
														GD,			
														LC,			
		LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NO,	NZ,	PL,	PT,	RO,
														UA,			
						AM,											
	RW:	GH,	GM,	KE,	LS,	MW,	ΜZ,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
		DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,
		ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG		
PRIORI'	TY APP	LN.	INFO	.:					US 2	000-	1956	22	P	2000	0406		
									US 2	001-	7660	53	A2	2001	0119		

OTHER SOURCE(S):

MARPAT 135:304105

GI

$$R^{6}$$
 R^{7}
 $N-R^{1}$
 R^{5}
 R^{4}
 R^{3}
 R^{2}
 I

AB Substituted nucleosides and isoindolinone derivs. I wherein Y is O, OH, S, Se, NH, N-alkyl, N-aryl; R1 is H, OH, aryl, alkyl, amino acid; R2 and R3 are independently H, alkyl, aryl, heterocycle, OH, O-alkyl, O-aryl, N-alkyl, N-aryl, taken together O, NH, S; R4-R7 are independently H, halo, alkyl-halo, OH, alkoxy, alkyl, alkenyl, carbocyclic, aryl, amino, carboxy, ester, arylalkyl, nitro; R3R4 are heterocyclic, carbocyclic ring; were prepd. as anti-inflammatory agents. Thus, isoindolinone I (Y = O, R1-R3 = R5-R7 = H, R4 = NO2) was prepd. and tested in vitro for its anti-inflammatory activity (% inhibition = 12 .mu.M).

IT 366454-42-4P 366454-43-5P 366454-45-7P

RL: BAC (Biological activity or effector, except adverse); IMF (Industrial manufacture); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of nucleosides and isoindolinone derivs. as anti-inflammatory

agents)

366454-42-4 HCAPLUS

RN Adenosine, 5'-deoxy-2',3'-O-(1-methylethylidene)-5'-[[1-oxo-3-(4,5,6,7-CN tetrahydro-4-oxo-1H-indol-1-yl)propyl]amino]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

366454-43-5 HCAPLUS RN

Adenosine, 5'-deoxy-5'-[[1-oxo-3-(4,5,6,7-tetrahydro-4-oxo-1H-indol-1-CN yl)propyl]amino] - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

366454-45-7 HCAPLUS RN

1H-Indole-1-propanoic acid, 4,5,6,7-tetrahydro-4-oxo- (9CI) (CA INDEX CN NAME)

=> d ibib abs hitstr 2

L18 ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2000:878646 HCAPLUS

DOCUMENT NUMBER: 134:178428

TITLE: Synthesis of 4,5,6,7-tetrahydroindole derivatives

AUTHOR(S): Zav'yalov, S. I.; Dorofeeva, O. V.; Rumyantseva, E.

E.; Kulikova, L. B.; Ezhova, G. I.; Kravchenko, N. E.;

Zavozin, A. G.

CORPORATE SOURCE: Zelinsky Institute of Organic Chemistry, Russian

Academy of Sciences, Moscow, Russia

SOURCE: Pharm. Chem. J. (2000), 34(3), 130-131

CODEN: PCJOAU; ISSN: 0091-150X
PUBLISHER: Consultants Bureau

DOCUMENT TYPE: Journal LANGUAGE: English

AB The condensation of 3-hydroxy-5,5-dimethyl-2-(2-oxo-2-phenylethyl)-2-cyclohexen-1-one with glycine gave 4,5,6,7-tetrahydro-6,6-dimethyl-4-oxo-2-phenyl-1H-indole-1-acetic acid. The condensation of the same starting

material with 4-aminobenzenesulfonamide gave (4,5,6,7-tetrahydro-6,6-dimethyl-4-oxo-2-phenyl-1H-indol-1-yl)benzenesulfonamide. The

cyclocondensation of glycine with phthalic acid gave N-(phthaloyl)glycine.

IT 121626-22-0P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation)
(prepn. of 4,5,6,7-tetrahydro-4-oxoindole derivs.)

RN 121626-22-0 HCAPLUS

CN 1H-Indole-1-acetic acid, 4,5,6,7-tetrahydro-6,6-dimethyl-4-oxo-2-phenyl-(9CI) (CA INDEX NAME)

IT 326809-42-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (prepn. of 4,5,6,7-tetrahydro-4-oxoindole derivs.)

RN 326809-42-1 HCAPLUS

CN 1H-Indole-1-acetic acid, 2-(4-bromophenyl)-4,5,6,7-tetrahydro-6,6-dimethyl-4-oxo-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

6

REFERENCE(S):

- (1) Anon; 1995, 21, HCAPLUS(2) Dagher, C; J Het Chem 1982, V19(3), P645 HCAPLUS
- (3) Nagarajan, K; J Med Chem 1976, V19(4), P508 HCAPLUS
- (5) Ramadas, S; Indian \ddot{J} Chem 1979, V17B(3), P195 **HCAPLUS**
- (6) Zav'Yalov, S; Khim-Farm Zh 1998, V32(3), P41 **HCAPLUS**
- ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d ibib abs hitstr 3

L18 ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2002 ACS 1997:479331 HCAPLUS ACCESSION NUMBER:

127:99527 DOCUMENT NUMBER:

Oxidative hair dye compositions containing TITLE:

n-substituted 4-hydroxy indoline derivatives Terranova, Eric; Fadli, Aziz; Lagrange, Alain

INVENTOR(S): Oreal S. A., Fr.

PATENT ASSIGNEE(S): Eur. Pat. Appl., 19 pp. SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 780118	A1	19970625	EP 1996-402297	19961029
EP 780118	B1	-19971229		
R: DE, ES,	FR, GB	, IT		
FR 2742047	A 1	19970613	FR 1995-14372	19951206
FR 2742047	B1	19980116		
CN 1189820	Α	19980805	CN 1996-195180	19960626
ES 2113769	Т3	19980501	ES 1996-402297	19961029
JP 09183716	A2	19970715	JP 1996-325758	19961205
JP 2996625	. B2	20000111		
US 5755829	Α	19980526	US 1996-761756	19961205
US 6002018	Α	19991214	US 1998-14622	19980128
PRIORITY APPLN. INFO	. :		FR 1995-14372 A	19951206
			US 1996-761756 A3	19961205

MARPAT 127:99527 OTHER SOURCE(S):

Oxidative hair dye compns. contain n-substituted 4-hydroxy indoline derivs. (Markush structure given). A soln. of 8.85 g 4-hydroxy-1-N-(.beta.-hydroxyethyl)indole (prepn. given) in 40 mL of acetic acid was stirred with 1.9 g of sodium cyanoborohydride at 30.degree. for 30 min, then the reaction mixt. was poured into 200 g water at pH = 7.5 and the ppt. thus obtained was filtered and dried to obtain 7.75 g 4-hydroxy-1-N-(.beta.-hydroxyethyl)indoline (I). A hair dye prepn. contained I 0.895, paraphenylenediamine 0.540, water and excipient q.s. 100 g. The hair dye prepn. is mixed with equal amt. of 20 vol. hydrogen peroxide and applied to the hair.

ΙT 186963-75-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (oxidative hair dye compns. contg. n-substituted 4-hydroxy indoline derivs.)

186963-75-7 HCAPLUS RN

4H-Indol-4-one, 1-(2,3-dihydroxypropyl)-1,5,6,7-tetrahydro- (9CI) CN INDEX NAME)

=> d ibib abs hitstr 4

L18 ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2002 ACS.

ACCESSION NUMBER:

1997:168570 HCAPLUS

DOCUMENT NUMBER:

126:185981

TITLE:

Preparation of N-hydroxyalkyl-4-hydroxyindoles as

oxidative hair dye components

INVENTOR(S):

Terranova, Eric; Fadli, Aziz; Lagrange, Alain

US 1996-678981

A3 19960712

PATENT ASSIGNEE(S):

L'Oreal S. A., Fr. Eur. Pat. Appl., 20 pp.

SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

French

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

PA'	TENT NO.	KIN	D DATE	APPLICATION NO.	DATE
	754681 754681	A1 B1	19970122 19980304		19960626
FR			DE, ES, FR, 19970117	GB, IT, LI, NL, SE FR 1995-8566	19950713
CA	2222312 9703049	AA A1	19970130 19970130	CA 1996-2222312 WO 1996-FR996	19960626 19960626
ES JP BR	W: BR, 163640 2117474 10512282 9609329 3095419	E T3	19981124 19990525	AT 1996-401413 ES 1996-401413 JP 1996-505543 BR 1996-9329	19960626 19960626 19960626 19960626
US US	5704948 5869692 Y APPLN.	A A INFO.:	19980106 19990209	US 1996-678981	19960712 19970918

OTHER SOURCE(S):

MARPAT 126:185981

GI

AB Title compds. (I; R1 = hydroxyalkyl, alkoxyalkyl, aminoalkyl, etc.; R2,R3 = H, halo, alkyl, CO2H, alkoxycarbonyl, CHO; R4 = H, halo, alkyl, alkoxy, etc.) were prepd. Thus, 4-oxo-4,5,6,7-tetrahydrobenzofuran was cyclocondensed with H2NCH2CH2OH and the product dehydrogenated to give I (R1 = CH2CH2OH, R2-R4 = H). Data for activity of I were given.

IT 186963-75-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. of N-hydroxyalkyl-4-hydroxyindoles as oxidative hair dye components)

RN 186963-75-7 HCAPLUS

CN 4H-Indol-4-one, 1-(2,3-dihydroxypropyl)-1,5,6,7-tetrahydro- (9CI) (CA INDEX NAME)

=> d ibib abs hitstr 5

L18 ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1989:457475 HCAPLUS

DOCUMENT NUMBER:

111:57475

TITLE:

Synthesis and oral hypoglycemic properties of 4-oxo-4,5,6,7-tetrahydroindole-3-acetic acids

AUTHOR(S):

Nagarajan, Kuppuswamy; Talwalker, Purnachand K.; Goud, A. Nagana; Shah, Rashmi K.; Shenoy, Sharada J.; Desai,

Narasimha D.

CORPORATE SOURCE:

Res. Cent., Hindustan Ciba-Geigy Ltd., Bombay, 400

063, India

SOURCE:

Indian J. Chem., Sect. B (1988), 27B(12), 1113-23

CODEN: IJSBDB; ISSN: 0376-4699

DOCUMENT TYPE:

PE: Journal

LANGUAGE: OTHER SOURCE(S): English CASREACT 111:57475

GΙ

Me OH CH
$$_{2}CO_{2}H$$

Me Me N $_{R}^{2}$

N $_{N}R^{3}$

PhN III

Condensation of .beta.-acetyl-2-hydroxy-4,4-dimethyl-6-oxo-1-cyclohexene-1-propionic acid (I) with NH4OAc and primary amines affords tetrahydroindole-3-acetic acids II (R = alkyl, aryl, aralkyl; R1 = Me, R2 = CH2CO2H), while another dimedone deriv. serves as starting material for isomeric indole-2-acetic acids II (R = alkyl, R1 = CH2CO2H, R2 = H). 4-Oxotetrahydroindole-2-carboxylic acids II (R = Ph, CH2CHMe2, R1 = CO2H, R2 = Me) and 3-carboxylic acids II (R = Ph, 4-FC6H4, R1 = H, R2 = CO2H), are obtained from the corresponding benzofurans. Some of the 3-carboxylic acid esters are transformed to tricyclic compds. like III [R3 = H, Me2N(CH2)3, Et2NCH2CH2]. Good oral hypoglycemic activity in normal rats is shown generally by the 3-acetic acids, among which C 8778-GO and C 9001-GO (II, R = Bu, CH2CHMe2, R1 = Me, R2 = CH2CO2H) are most active. These two acids are also active in streptozotocin-induced diabetic rats and have been investigated extensively. Structure-activity relationships are discussed.

IT 121625-91-0P 121626-21-9P 121626-22-0P

121626-23-1P 121626-54-8P

RL: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prepn. and hypoglycemic activity of)

RN 121625-91-0 HCAPLUS

CN 1H-Indole-1-butanoic acid, 3-(carboxymethyl)-4,5,6,7-tetrahydro-2,6,6-trimethyl-4-oxo- (9CI) (CA INDEX NAME)

RN 121626-21-9 HCAPLUS CN 1H-Indole-1-acetic acid, 4,5,6,7-tetrahydro-2,6,6-trimethyl-4-oxo- (9CI) (CA INDEX NAME)

RN 121626-22-0 HCAPLUS CN 1H-Indole-1-acetic acid, 4,5,6,7-tetrahydro-6,6-dimethyl-4-oxo-2-phenyl-(9CI) (CA INDEX NAME)

RN 121626-23-1 HCAPLUS CN 1H-Indole-1-butanoic acid, 4,5,6,7-tetrahydro-2,6,6-trimethyl-4-oxo- (9CI) (CA INDEX NAME)

RN 121626-54-8 HCAPLUS CN 1H-Indole-1-butanoic acid, 2-(carboxymethyl)-4,5,6,7-tetrahydro-6,6dimethyl-4-oxo- (9CI) (CA INDEX NAME)